PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6: A61K 9/14, 9/72, 47/12, B01J 2/28

(11) International Publication Number:

WO 95/05805

(43) International Publication Date:

2 March 1995 (02.03.95)

(21) International Application Number:

PCT/SE94/00780

(22) International Filing Date:

25 August 1994 (25.08.94)

(30) Priority Data:

9302777-9

27 August 1993 (27.08.93)

SE

(71) Applicant (for all designated States except US): ASTRA AKTIEBOLAG [SE/SE]; S-151 85 Södertälje (SE).

(72) Inventors; and

(75) Inventors/Applicants (for US only): TROFAST, Eva, Ann-Christin [SE/SE]; Vapenkroken 34, S-226 47 Lund (SE). BRIGGNER, Lars-Erik [SE/SE]; Arkeologvägen 27, S-226 54 Lund (SE).

(74) Agent: ASTRA AKTIEBOLAG; S-151 85 Södertälje (SE)

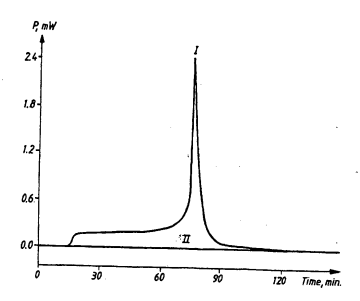
(81) Designated States: AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, JP, KE, KG, KP, KR, KZ, LK, LT, LU, LV, MD, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SI, SK, TI, TT, UA, US, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG), ARIPO patent (KE, MW, SD).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: PROCESS FOR CONDITIONING SUBSTANCES



(57) Abstract

The present invention relates to a process for providing a stable crystallinic form to a fine-grained substance or a substance mixture, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such a substance or a substance mixture, by a) in case of a substance mixture, preparing a homogeneous mixture of the substances; b) micronizing, direct precipitating or diminishing by any conventional method the substance or substance mixture into a particle size required for inhalation, the particle size being less than $10 \mu m$; c) optionally preparing a homogeneous mixture of the desired substances when each substance has been introduced from stage b) as separate fine-grained particles; d) conditioning said substance or substance mixture by treatment with a water containing vapour phase in a controlled fashion; and e) drying.

3

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT Australa AU Australia BB Barbados GE Georgia BB Gelgrium GR Greece BF Burkina Faso HU Hungary BJ Benin BR Brazil BY Belarus KE Kenya CA Canada KG Kyrgystan CF Central African Republic CG Congo CH Switzerland KR Republic of Korea CH Cameroon LI Liecternstein CN Cana CN Canad LK Sri Lanka CN Chaa CC Czechoslovakia LU Luxembourg CE Germany MC Monaco FI Lizekinan MR Mauritania MW Malawi MR Mauritania MW Malawi Malaw						
BB Barbados GN Guinea NE Niger BE Belgium GR Greece NL Netherlands BF Burkina Faso HU Hingsry NO Norway BJ Benin IT Italy PL Poland BR Brazil JP Japan PT Portugal BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KF Democratic People's Republic SD Sudan CH Switzerland KR Republic of Korea SE Sweden CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germany MC Monaco TT Trinidad and Tobago EN Gameroen MG Madagasear US United States of America FI Finland ML Mali UZ Uzbekistan			GB	United Kingdom	MD	Marwisonia
BE Belgium GR Greece NL Netherlands BF Burkina Faso HU Hungary NO Norway BG Bulgaria IE Ireland NZ New Zealand BR Brazil JP Japan PT Portugal BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KP Democratic People's Republic SD Sudan CH Switzerland KR Republic of Korea SI Slovenia CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CZ Czechoslovakia LU Luxembourg TG Togo DE Germany MC Monaco TT Trinidad and Tobago DK Demmark MD Republic of Moldova UA Utraine FR Finland ML Mali UZ Uzbekistan			GE	Georgia		
BF Burkina Faso HU Hungary NO Norway BG Bulgaria IE Ireland NZ New Zealand BR Brazil JP Japan PT Poland BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KP Democratic People's Republic SD Sudan CH Switzerland KR Republic of Korea SE Sweden CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CZ Czechoslovakia LU Luxembourg TG Togo DE Germany MC Monaco TT Trinidad and Tobago DK Demark MD Republic of Moldova UA Ukraine FR France MN Monaco III Uzbekistan MG Madagascar US Uzbekistan			GN	Guinea		
BY Burkins Faso BI Bulgaria BG Bulgaria BI Benin BR Brazil BP Japan BY Belarus CA Canada CF Central African Republic CG Congo CH Switzerland CH Switzerland CM Cameroon LI Liechtenstein CM Cameroon LI Liechtenstein CN China CN China CN China CN China CN Czechoslovakia CN Czechoslovakia CN Czechoslovakia CN Cermany CN MC Monaco CM Monaco CM Cameron CR Camero CR Came		_	GR	Greece		
BUJ Benin II Italy PL Poland BR Brazil JP Japan PT Portugal BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SI Slovenia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CC Czech Republic LV Latvia TJ Tajikistan CC Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germany MC Monaco TT Trinidad and Tobago DE DE Germark MD Republic of Moldova UA Ukraine FI Finland ML Maij UZ Uzbekistan FR France MN Monaco NZ New Zealand NEW Jelland NZ New Zealand NZ New Zealand NEW Jelland NZ New Zealand NEW Jelland NZ New Zealand NE Welland NZ New Zealand NEW Jelland NZ New Zealand NEW Jelland NZ New Zealand NEW Jelland NZ New Zealand NEW Jelland NEW Jelland NZ New Zealand NEW Jelland NEW Jelland NZ New Zealand NZ New Zealand NZ New Zealand NZ New		Burkina Faso	EU			
BJ Benin IT Italy PL Poland BR Brazil JP Japan PT Portugal BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KP Democratic People's Republic SD Sudan CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SI Slovenia CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CC Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germark MD Republic of Moldova UA Ukraine ES Spain MG Madagasear US United States of America FR Finland ML Mali UZ Uzbekistan	_	Bulgaria				
BR Brazil JP Japan PT Pottugal BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KP Democratic People's Republic SD Sudan CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SI Slovenia CI Côte d'Ivoire KZ Kazakhstan SI Slovenia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxenbourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan CK Germany MC Monaco TT Trinidad and Tobago DE Germany MC Monaco TT Trinidad and Tobago DE Demark MD Republic of Moldova UA Ukraine FI Finland ML Mali UZ Uzbekistan MN Monacoi UZ Uzbekistan	BJ	Ben <u>in</u>	Fr		_	
BY Belarus KE Kenya RO Romania CA Canada KG Kyrgystan RU Russian Federation CF Central African Republic KP Democratic People's Republic SD Sudan CH Switzerland KR Republic of Korea SE Sweden CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CZ Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Demark MD Republic of Moldova UA Utraine FI Finland ML Mali UZ Uzbekistan FR France MN Monaco UZ Uzbekistan	BR	Brazil		•		
CA Canada KG Kyrgystan RU Romania CF Central African Republic KP Democratic People's Republic SD Sudan CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SI Slovenia CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germany MC Monaco TT Trinidad and Tobago DE Demark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan	BY	Belerus	-	•		
CF Central African Republic KP Democratic People's Republic SD Sudam CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SI Slovenia CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan	CA	Canada				
CG Congo of Korea SE Sweden CH Switzerland KR Republic of Korea SE Sweden CI Côte d'Ivoire KZ Kazakhstan SK Slovakia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CZ Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TJ Tajikistan DK Demmark MD Republic of Moldova UA Ukraine ES Spain MG Madagasear US United States of America FI Finland ML Mali UZ Uzbekistan	CF	Central African Republic	_			Russian Federation
CH Switzerland KR Republic of Korea SI Shovenia CI Côte d'Ivoire KZ Kazakhstan SI Slovenia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TJ Trinidad and Tobago DK Demark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan	CG		IS.F	Democratic People's Republic		Suden
CI Côte d'Ivoire KZ Kazakhstan SI Slovenia CM Cameroon LI Liechtenstein SN Senegal CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TT Trinidad and Tobago DE Germark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan	CH	_	Ph.	-	SE	Sweden
CM Cameroon LI Liechtenstein SK Slovakia CN China LK Sri Lanka TD Chad CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TJ Trinidad and Tobago DE Demark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan	CI		-		SI	Slovenia
CN China LL Lechtenstein SN Senegal CS Czechoslovakia LL Luxembourg TG Togo CZ Czech Republic LV Luxem TT Tajikistan DE Germany MC Monaco TJ Tajikistan DK Demmark MD Republic of Moldova UA Ukraine ES Spain MG Madagasear US United States of America FR France MN Monaco UZ Uzbekistan	CM				SK	Slovakia
CS Czechoslovakia LU Luxembourg TG Togo CZ Czech Republic LV Latvia TJ Tajikistan DE Germany MC Monaco TJ Tajikistan DK Dermark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Maji UZ Uzbekistan	CN				SN	Senegal
CZ Czech Republic LV Latvia TJ Tajikiritan DE Germany MC Monaco TJ Trinidad and Tobago DK Demmark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Maji UZ Uzbekistan					TD	
DE Germany MC Monaco TJ Tajikistan DK Demark MD Republic of Moldova UA Ukraine ES Spain MG Madagascar US United States of America FI Finland ML Mali UZ Uzbekistan					TG	Togo
DK Demark MC Monaco TT Trinidad and Tobago ES Spain MG Madagascar UA Ukraine FI Finland ML Mali US United States of America FR France MN Monachis					TJ	•
ES Spain MD Republic of Moldova UA Ukraine FI Finland ML Mali UZ Uzbekistan FR France MN Monorbis	-				Tr	
FI Finland ML Maji UZ Uzbekistan MG Madagascar US United States of America FR France MN Managalia UZ Uzbekistan				Republic of Moldova		
FR France MN Mongolia UZ Uzbekistan		•	MG	Madagascar		
MN Monardia			ML	Maij		
GA Gabon VII VIC Nami		-	MN	Mongolia		
	GA	Gabon		-	*14	A NOT TAKEN

Process for conditioning substances

5

Field of the invention

The present invention relates to a process for providing a fine-grained substance or substance

mixture, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such a substance or substance mixture and which have improved physicochemical properties in the dry state, thereby facilitating the technical handling and significantly increase the medical value of the formulation used.

Background of the invention

- There are presently several effective drugs available for the treatment of patients with asthma or other respiratory disorders. It has been recognized that these drugs should be given by the inhaled route whenever possible. The ideal delivery system for inhalable drugs would be a user- and environment-friendly multidose inhaler giving accurate doses of a stable formulation with good aerodynamic behaviour of the particles.
- During the past few years, there have been frequent demonstrations of the fact that the appropriate selection of the most suitable crystalline modification significantly can influence the clinical results of a given chemical substance. The chemical and physical stability of a solid in a particular dosage form can be
- improved by presenting the substance(s) in the appropriate crystal form. The solid state phase

transformation of the substance in a dosage form can dramatically alter the pharmaceutical properties of the formulation. The solid state phase of the administered substance(s) can influence such important factors as bioavailability and physicochemical stability (specific surface area, particle size etc). Chemical stability in solid state and hygroscopicity are often closely related to the crystallinity.

- Solid state transformations may occur during mechanical processing e.g. micronization. In a micronization process of solids, disruption or activation of the crystalline structure often leads to varying degrees of disorder through the formation of defects or amorphous
- regions. Such regions are often more sensitive to external effects e.g. moisture. It is necessary to establish the conditions whereby different forms of a substance might be converted to a single stable form thus eliminating differences in solid state properties and subsequent different physicoghomical and
- and subsequent different physicochemical and pharmaceutical properties.

The increasing production and use of fine powders in the pharmaceutical industry has highlighted the need of reliable methods for assessing their physicochemical 25 and technical handling. Mixing of cohesive powders will be influenced by the interparticulate forces between particles of the same species and also between particles of different species. Since fine powders agglomerate, the mixture will often be inhomogeneous, 30 particularly the minor component will show a skewed distribution. One reason could be that the agglomerates of the minor component is not completely dispersed into their component particles; see further Chem. Eng. (1973), 12-19. Cohesive powders are thus very difficult 35 to mix to a homogenous mixture in an accurate way,

esp cially when one component is present only as a

small fraction.

Substances will often be obtained in an amorphous state or a metastable crystalline form when spray drying, freeze drying, rapid solvent quenching or when using 5 controlled precipitation, where both crystalline and amorphous forms can be prepared. The use of an amorphous form or metastable crystalline form is often limited due to its thermodynamic instability. It is therefore a desire to convert the amorphous form or the 10 metastable crystalline form to the more stable crystalline state. For crystalline substances, a diminution operation step will give amorphous regions of the particle making the particle more sensitive to moisture and chemical degradation. The present 15 invention deals with such physical changes, or more importantly, to anticipate them and the means by which

The rearrangement or conditioning of a water-soluble substance, amorphous or partly amorphous, using a solvent like ethanol, acetone or the like has been described in Eur. Pat. Appl. EP 508 969 where single compounds have been applied. However, that method is not applicable for some substances containing crystal water, since organic solvents will eliminate the water thereby changing the properties of the substance considerably. It has been understood that water-soluble substances could not be conditioned by water while

keeping the particle distribution of a fine-grained

these solid state phenomena can be handled.

References:

substance intact.

30

Amorphous-to-Crystalline Transformation of Sucrose,

Phar. Res., 7(12), 1278 (1990) by J.T. Carstensen and
K. Van Scoik.

Effect of Surface Characteristcs of Theophylline

15

20

Anhydrate Powder on Hygroscopic Stability, J. Pharm. Pharmacol. 42, 606 (1990) by M. Otsuka et al. Process for conditioning of water-soluble substances, Eur. Pat. Appl. 508969 by J. Trofast et al.

The molecular basis of moisture effect on the physical and chemical stability of drugs in the solid state, Int. J. Pharm. 62(1990), 87-95 by C. Ahlneck and G. Zografi.

10 Brief description of the invention

The object of the invention is to provide a process for a fine-grained substance or substance mixture, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such a substance or substance mixture, whereby conditioning the mixture in a controlled process, thereby facilitating the technical handling and significantly increase the medical value of the formulation used.

Detailed description of the invention

The object of the present invention is to provide a reliable process for providing a stable crystallinic form to a fine-grained substance or a substance mixture, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such a substance or a substance mixture. The process according to the present invention comprises the following steps:

- a) in case of a substance mixture, preparing a homogenous mixture of the substances;
- b) micronizing, direct precipitating or diminishing by any conventional method the substance or substance mixture into a particle

5

size required for inhalation, the particle size being less than $10\mu m$;

- c) optionally preparing a homogenous mixture of the desired substances when each substance has been introduced from stage b) as separate fine-grained particles;
- d) conditioning said substance or substance
 mixture by treatment with a water containing
 vapour phase in a controlled fashion; and
 - e) drying.

transition).

- The conditioning step is carried out by treatment with a water containing vapour phase. Said water containing vapour phase is a water vapour phase with or without any organic solvent vapour present.
- The conditioning step is carried out at a temperature/relative humidity combination, which suppresses the glass temperature of substances involved below the process temperature. The glass temperature (T_g) is the temperature at which the mobility of an amorphous material undergoes changes from an immobile glassy state to mobile rubbery state (phase
- The conditioning is generally carried out at a

 temperature between 0 and 100°C, preferably between 10
 and 50°C. Of practical reasons the conditioning is
 often performed at ambient temperature. The relative
 humidity (RH) at which the conditioning is carried out
 is chosen so that the phase transition occurs, mainly
 above 35% RH, preferably above 50% RH, and most
 preferably above 75% RH. The time used is considerably
 influenced by the batch size, relative humidity and

packing etc and may be from minutes to days.

The final formulation may also include different additives, e.g. a substance which enhances the absorption of a pharmacologically active drug in the lung. The enhancers used can be any of a number of compounds which act to enhance absorption through the layer of epithelial cell lining the alveoli of the lung and into the adjacent pulmonary vasculature. Among the substances with known absorption-enhancing properties are surfactants, such as alkali salts of fatty acids, sodium tauro-dihydrofusidate, lecithins, sodium glycocholate, sodium taurocholate, octylglucopyranoside and the like.

15

10

5

Other additives in the formulation may be carriers, diluents, antioxidants, buffer salts and the like, all of which will be treated according to the process of the present invention.

20

25

30

35

The accuracy and reproducibility of doses are often not sufficient when using very small doses in an inhalation device. Therefore very potent drugs may be diluted with a carrier in order to get an amount of powder sufficient to obtain a reliable and reproducible dose. Such a carrier may be carbohydrates like lactose, glucose, fructose, galactose, trehalose, sucrose, maltose, raffinose, maltitol, melezitose, starch, xylitol, mannitol, myoinositol, and the like and its hydrates, preferably lactose and mannitol, and amino acids such as alanine, betaine and the like.

Coarser particles having a size above 10 μm may also be conditioned using the process according to the present invention.

5

10

The present invention may be applied to for example the following pharmacologically active substances:

Formoterol (e.g. as fumarate) and salmeterol (e.g. as xinafoate) are highly selective long-acting β_2 -adrenergic agonists having bronchospasmolytic effect and are effective in the treatment of reversible obstructive lung ailments of various genesis, particularly asthmatic conditions. Salbutamol (e.g. as sulphate), bambuterol (e.g. as hydrochloride), terbutaline (e.g. as sulphate), fenoterol (e.g. as hydrochloride), procaterol (e.g. as hydrochloride), bitolterol (e.g. as hydrochloride), procaterol (e.g. as hydrochloride), bitolterol (e.g. as mesylate) and broxaterol are highly selective β_2 -

- adrenergic agonists and ipratropium bromide is an anticholinergic bronchodilator. Examples on antiinflammatory glucocorticoids are budesonide, (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-4-pregnen-3,20-dione, fluticasone
- (e.g. as propionate ester), beclomethasone (e.g. as dipropionate ester), tipredane, momethasone and the like. Several of the compounds could be in the form of pharmacologically acceptable esters, salts, solvates, such as hydrates, or solvates of such esters or salts, if any.

The preferred substances to which the invention is to be applied are terbutaline sulphate, salbutamol sulphate, fenoterol hydrobromide, ipratropium bromide, bambuterol hydrochloride, formoterol fumarate and salmeterol xinafoate, and their solvates, especially their hydrates.

The most preferred substance mixture to which the
invention is to be applied is formoterol (as formoterol
fumarate dihydrate)/lactose (monohydrate), although the
same principle may be applied to combinations such as

salbutamol (as salbutamol sulphate)/lactose, terbutaline (as terbutaline sulphate)/lactose, ipratropium bromide/lactose, budesonide/lactose, (22R)-6 α ,9 α -difluoro-11 β ,21-dihydroxy-16 α ,17 α -

- propylmethylenedioxy-4-pregnen-3,20-dione/mannitol,
 (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17αpropylmethylenedioxy-4-pregnen-3,20-dione/myoinositol
 and (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17αpropylmethylenedioxy-4-pregnen-3,20-dione/lactose. When
- one of the components is rather insoluble in water, it is possible to use an organic solvent as a conditioning agent for one compound and water vapour as a conditioning agent for the other one in the conditioning step. In that case the conditioning may be
- carried out in a two step procedure wherein the first step is conditioning with an organic solvent followed by conditioning by water vapour in a second step; or vice versa.
- The rearrangement or conditioning of the substance or substance mixture, amorphous or partly amorphous, involve treatment of the substance(s) with a water containing vapour phase in a controlled fashion. This conditioning step is to be performed in a defined environment with controlled and additional to the controlled and additi
- environment with controlled and adjustable humidity or a column using inert gas and/or organic solvent vapour containing the required amount of water vapour. The packing of the substance or substance mixture affects the time needed as well as the result of the
- conditioning. The tendency of caking is affecting the number and size of particles. In case of a substance mixture, it is usually an advantage to mix the substances before the micronizing step in order to ensure a homogenous mixture when using small ratios
- 35 between the drug substance and the additive.

With the present inv ntion it is possible to condition two or more substances in the same process while the particle distribution is maintained and this is from a technical standpoint a great advantage.

5

10

15

20

The ratio between the substances in a substance mixture is between 1:1 and 1:1000, preferably between 1:1 and 1:500, and most preferred between 1:1 and 1:200 in the case where one substance is a pharmacologically active substance and the other one is an additive.

The particle size of the fine-grained substances should be identical before and after the conditioning step as measured by different instruments like Malvern Master Sizer, Coulter Counter or a microscope.

It is also of utmost importance that the particles obtained are well-defined in size and distribution as well as have small batch to batch variations in order to obtain agglomerates that will completely disintegrate into its primary particles in the inhaler used.

It is an object of the present invention to provide a reliable process, where the drug formulation of a single drug substance or a combination of a drug substance/additive, preferably formoterol fumarate dihydrate/lactose can be conveniently and reproducibly prepared.

30

35

For some material such as formoterol/lactose, where the T_g (the glass transition temperature, the temperature at which the mobility of an amorphous substance undergoes changes from an immobile glassy state to mobile rubbery state) or water sensitivity is markedly different for the drug substance and the additive, the process can be performed in two subsequent steps, i.e.

conditioning of one substance at one temperature/RH combination followed by conditioning at a higher temperature/RH for a second substance.

The mixing step is preferably performed before the micronization step in order to ensure the content uniformity or in a single step using a vibratory ball mill as reported by I. Krycer and J.A. Hersey in Int. J. Pharm. 6, 119-129 (1980). It is also possible to mix the substances after micronization or after each substance has been conditioned.

In some instances it has been possible to use infrared spectroscopy in order to study the conversion of an amorphous form or a partly crystalline form, into a stable crystalline form. Other methods available include BET gas adsorption, X-ray powder diffraction, isothermal microcalorimetry and differential scanning calorimetry (DSC). We have found that BET gas adsorption and isothermal microcalorimetry being the best methods for distinguishing the different forms of the tested compounds.

When a substance or substance mixture is agglomerated and used as such, a drop of about 70-80% of the 25 respirable particles is found when exposed to high humidity. It has astonishly been found that a drop of only about 25-30% occurs when a substance or substance mixture has been conditioned (at 50% RH for formoterol fumarate dihydrate/lactose mixture) before 30 agglomeration and exposed to high humidity. After further conditioning at 75% RH a drop of only 5-10% of the respirable particles will occur. There is no difference in particle distribution as measured by a Malver instrument before and after conditioning at 75% 35 RH. If the conditioning is performed with the agglomerated product the particle distribution is

considerable worse and the formulation useless in an inhalation device.

Experimental procedure

5

The invention relates to the following procedure:

1. Mixing the drug substance with the additive in a defined ratio.

10

- 2. Micronizing the mixture.
- 3. Conditioning at a temperature/relative humidity combination, which suppresses the glass temperature of substances involved below the process temperature. The glass temperature (T_g) is the temperature at which the mobility of an amorphous material undergoes changes from an immobile glassy state to mobile rubbery state.
- 20 4. Drying with dry nitrogen or air, or in vacuum.

EXAMPLES

The invention is further illustrated but not limited by the following examples performed according to the described experimental procedure. Several batches of each substance or substance mixture have been measured. The data represents a comparison of the heat (J/g) given off by non-conditioned and conditioned substances when subjected to a water containing vapour phase. The experiments are performed by using a Thermal Activity Monitor 2277 (Thermometrics AB, Sweden).

Example 1

Salbutamol sulphate (25%)/lactose (75%)

Conditioned at relative humidity (RH)

Non-conditioned substance (J/g)

Conditioned substance (J/g)

50-60 % RH

5-8

Conditioned substance (J/g)

<0.5

	Example 2	
	Ipratropium bromide (6%)/lactose (94%)	
·	Conditioned at relative humidity (RH)	50-60 % RH
5	Non-conditioned substance (J/g)	6-8
	Conditioned substance (J/g)	<0.5
	Example 3	
	Formoterol fumarate dihydrate	
10	Conditioned at relative humidity (RH)	75 % RH
	Non-conditioned substance (J/g)	6
	Conditioned substance (J/g)	<0.5
15	Example 4	·
	<u>Lactose (see Figure 1)</u>	
	Conditioned at relative humidity (RH)	50 % RH
	Non-conditioned substance (J/g)	10-14
20	Conditioned substance (J/g)	<0.5
	Example 5	
•	Melezitose	
•	Conditioned at relative humidity (RH)	50 % RH
25	Non-conditioned substance (J/g)	12
•	Conditioned substance (J/g)	<0.5
	Example 6	
	Formoterol fumarate dihydrate (2%)/lactose	e (98%)
30	Conditioned at relative humidity (RH)	50 % RH
•	Non-conditioned substance (J/g)	10-14
	Conditioned substance (J/g)	<0.5
	During a recrystallization a large amount	of heat is
35	evolv d, and by monitoring the calometrical	or mean to
	sample is checked for any amorphous content	t. Figure 1
	shows micronised lactose before (I) and af	ter (II)

conditioning. Thus, a complete crystallinity has been obtained during the conditioning according to the invention.

5

CLAIMS

- 1. A process for providing a stable crystallinic form to a fine-grained substance or a substance mixture, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such a substance or a substance mixture,
 10 characterized in
 - a) in case of a substance mixture, preparing a homogenous mixture of the substances;
- b) micronizing, direct precipitating or diminishing by any conventional method the substance or substance mixture into a particle size required for inhalation, the particle size being less than 10 µm;
 - c) optionally preparing a homogenous mixture of the desired substances when each substance has been introduced from stage b) as separate fine-grained particles;
 - d) conditioning said substance or substance mixture by treatment with a water containing vapour phase in a controlled fashion; and
- e) drying.

20

25

A process according to claim 1 characterized in that the conditioning, in the case of
a substances mixture, may be performed in a one step
procedure or a multistep procedure using different
relative humidity/temperature combinations.

30

35

- 3. A process according to claim 1
 charact rized in that the substance or substance mixture is a drug formulation of a single drug substance or a combination of a drug substance/additive.
- 4. A process according to claim 1
 c h a r a c t e r i z e d that said substance or at least one of the substances of said substance mixture
 is selected from formoterol, salmeterol, salbutamol, bambuterol, terbutaline, fenoterol, clenbuterol, procaterol, bitolterol, broxaterol, ipratropium bromide, budesonide, (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-4-pregnen-3,20-dione, fluticasone, beclerathy.
- dione, fluticasone, beclomethasone, tipredane, momethasone, and pharmacologically acceptable esters, salts, solvates, such as hydrates, and solvates of such esters or salts, if any.
- 5. A process according to claim 1
 c h a r a c t e r i z e d in that said substance or at least one of the substances of said substance mixture is selected from formoterol fumarate, salmeterol xinafoate, salbutamol sulphate, bambuterol
- hydrochloride, terbutaline sulphate, fenoterol hydrobromide, clenbuterol hydrochloride, procaterol hydrochloride, bitolterol mesylate, fluticasone propionate, beclomethasone dipropionate and solvates, such as hydrates thereof, if any.
 - 6. A process according to claim 3

 c h a r a c t e r i z e d that the additive is a carrier selected from lactose, glucose, fructose, galactose, trehalos, sucrose, maltose, raffinose, maltitol, melezitos, starch, xylitol, mannitol, myoinositol, and the like, and its hydrates, preferably lactose and mannitol, and amino acids such as alanine,

betaine and the like.

7. A process according to claim 3

c h a r a c t e r i z e d that the additive is an enhancer selected from surfactants, such as alkali salts of fatty acids, sodium tauro-dihydrofusidate, lecithins, sodium glycocholate, sodium taurocholate, octylglucopyranoside and the like, or an antioxidant or a buffer salt.

10

5

- 8. A process according to claim 1
 c h a r a c t e r i z e d in that said substance mixture is selected from formoterol/lactose,
 salbutamol/lactose, terbutaline/lactose, ipratropium
 bromide/lactose, budesonide/lactose, (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-4-pregnen-3,20-dione/mannitol, (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-4-pregnen-3,20-dione/myoinositol and (22R)-6α,9α-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-difluoro-11β,21-dihydroxy-16α,17α-propylmethylenedioxy-
- 9. A process according to claim 1
 c h a r a c t e r i z e d in that said substance
 mixture is selected from formoterol fumarate
 dihydrate/lactose, salbutamol sulphate/lactose and
 terbutaline sulphate/lactose.
 - 10. A process according to claim 1

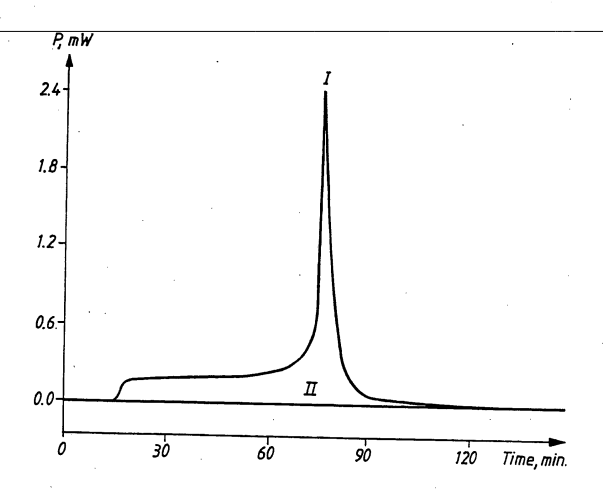
4-pregnen-3,20-dione/lactose.

30 characterized that step d) is carried out at a temperature between 0 and 100°C, preferably between 10 and 50°C and at a relative humudity so as that the phase transition occurs, mainly above 35% RH, preferably above 50% RH, and most preferably above 75% RH.

11. A process according to claim 1
c h a r a c t e r i z e d in that the ratio between
the substances in a substance mixture is between 1:1
and 1:1000, preferably between 1:1 and 1:500, and most
preferred between 1:1 and 1:200 in the case where one
substance is a pharmacologically active substance and
the other one is an additive.

1/1

F-19.1



INTERNATIONAL SEARCH REPORT

International application No. PCT/SE 94/00780

A. CLASSIFICATION OF SUBJECT MATTER		
IPC6: A61K 9/14, A61K 9/72, A61K 47/12, According to International Patent Classification (IPC) or to both	B01J 2/28 national classification and IPC	
B. FIELDS SEARCHED Minimum documentation searched (classification system followed in the searched)	hu ala sei Casai ann an 1919	
William Goodine Marion searched (Classification system followed	by classification symbols)	
IPC6: A61K, B01J		
Documentation searched other than minimum documentation to the	ne extent that such documents are included i	n the fields searched
SE,DK,FI,NO classes as above		
Electronic data base consulted during the international search (nam	e of data base and, where practicable, search	h terms used)
		•
WPI, WPIL, CLAIMS, EMBASE, CHEMICAL ABS	TDACT	
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where ap	opropriate, of the relevant passages	Relevant to claim No.
A EP, A1, 0508969 (AKTIEBOLAGET A	STRA),	1-11
14 October 1992 (14.10.92)		
		٠.
A WO, A1, 9116882 (LIPOSOME TECHNO	או חפע זאר ז	1-11
14 November 1991 (14.11.91)	secon, inc.),	- 1-11
WO, A1, 8400294 (SCHRÖDER, ULF) (02.02.84)	, 2 February 1984	1-11
(02.02.04)	•	•
		
·		
Further documents are listed in the continuation of Bo	x C. X See patent family annex	
 Special categories of cited documents: "A" document defining the general state of the art which is not considered 	"T" later document published after the inte date and not in conflict with the application of the principle on the published after the inter-	cation but cited to understand
to be of particular relevance "E" entire document but published on or after the international filing date	"X" document of particular relevance: the	claimed invention cannot be
"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other	considered novel or cannot be conside step when the document is taken alone	red to involve an inventive
special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance: the considered to involve an inventive step	claimed invention cannot be
means "P" document published prior to the international filing date but later than	combined with one or more other such	documents, such combination
the priority date claimed	"&" document member of the same patent	family
Date of the actual completion of the international search	Date of mailing of the international s 1 9 -12- 1994	earch report
13 December 1994		
Name and mailing address of the ISA/ Swedish Patent Office	Authorized officer	
Box 5055, S-102 42 STOCKHOLM	 Anneli Jönsson	
Facsimile No. +46 8 666 02 86	Telephone No. +46 8 782 25 00	
orm PCT/ISA/210 (second sheet) (July 1992)		

INTERNATIONAL SEARCH REPORT

Information on patent family members

26/11/94

International application No.
PCT/SE 94/00780

Patent document cited in search report		Publication date	Patent family member(s)		Publication date
EP-A1-	0508969	14/10/92	AU-A- CA-A- CZ-A- EP-A- HU-A- HU-D- JP-T- WO-A-	1534792 2106975 9302116 0580648 65095 9302870 6506454 9218110	17/11/92 12/10/92 13/04/94 02/02/94 28/04/94 00/00/00 21/07/94 29/10/92
WO-A1-	9116882	14/11/91	AU-A- EP-A-	7908791 0527940	27/11/91 24/02/93
WO-A1-	8400294	02/02/84	AU-B- AU-A- DE-A- EP-A,B- US-A-	567434 1776083 3376797 0113749 4713249	19/11/87 08/02/84 07/07/88 25/07/84 15/12/87

Form PCT/ISA/210 (patent family annex) (July 1992)